

(1) Publication number:

0 161 328

CUDODEAN	DATENT	APPLICATION
FUROPEAN	PAIENI	APPLICATION

- Application number: 84105636.9
 - Date of filing: 17.05.84

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⑤ Int. Cl.*: G 01 N 33/569, G 01 N 33/546, C 12 N 7/06, C 12 N 7/02

- Date of publication of application: 21.11.85
 Bulletin 85/47
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- Supported viral antigen and preparation and use thereof.
- A solid support is sensitized with soluble rubella virus antigen which is obtained by disruption and solubilization of whole (intact) rubella virus. The sensitized support is useful in an assay for rubella virus antibody.

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Opposition against EP Patent 967484 / 98 93 5359.4 Our Ref.: N3240 EP/OPP S3

This invention relates to viruses, and more particularly 11 2 to the purification of virus, production of virus antigens, the use of virus antigens for the production of sensitized solids and the use of virus antigen sensitized solids for testing for virus antibodies. Most particularly, the invention relates to rubella virus, rubella virus antigen and a test for rubella virus 7 antibody. United States Patent No. 4,195,074 discloses a process

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for producing soluble rubella virus antigen, and the use thereof in an agglutination test for rubella virus antibody. In accordance with U.S. Patent 4,195,074, the tissue culture from rubella virus infected cells is subjected to immunosorbent separation through a column containing IgG derived from human serum known to contain antibodies reactive with rubella antii followed by elution of the rubella antigen material from the 15 column and selection of the soluble antigen by gel permeation 16 chromatography. The antigen may then be employed for sensitizing erythrocytes, and the sensitized erythrocytes are used to deter-18 mine antibody in human serum samples by direct agglutination. 19

In accordance with the aforesaid patent, the so-called rubella antigen is not recovered from the virus, per se, and, therefore, it is believed that such material does not include structural proteins of the virus.

In accordance with one aspect of the present invention, there is provided a solid support sensitized with soluble rubella viral antigen which is obtained by disruption and solubilization of whole intact rubella virus.

In accordance with another aspect of the invention, soluble rubella virus antigen is obtained from whole rubella virus.

In accordance with still another aspect of the present

invention, there is provided a test or assay for rubella virus antibody and a reagent kit therefor.

In accordance with a further aspect of the present invention, there is provided a process for producing purified virus by the use of an adsorption gel to remove non-viral proteins and nucleic acids.

In accordance with yet a further aspect of the invention, there is provided a method for producing a solid sensitized with a viral antigen.

More particularly, the rubella virus antigen is isolated from intact rubella virus by treating purified whole rubella virus with a surfactant or detergent which disrupts the virus to provide the soluble rubella virus antigen, without destroying the antigenic characteristics thereof. The detergent is employed in an amount that is sufficient to disrupt and solubilize the whole virus without destroying its antigenic characteristics.

The surfactant or detergent which is used for disrupting the whole rubella virus may be any one of a wide variety of surfactants or detergents which disrupt and solubilize the virus, without destroying the antigenic characteristics, including cationic, anionic and non-ionic surfactants. Such surfactants are well known in the art, and as representative examples, there may be mentioned alkali metal salts of sulfates, soaps, sulfated or sulfonated oils, various amines, quaternary salts, condensation products with ethylene oxide, etc. Such detergents and surfactants and the use thereof for disrupting whole virus are known in the art. Preferred detergents for such use are alkali (lithium or sodium) dodecyl sulfate, sulfobetain, deoxylcholate and laurolybsarcosine (Sarcosyl).

In the case where the rubella virus antigen is to be supported on a solid support for use in an agglutination assay technique, the detergent or surfactant is one which is capable 3 of disrupting and solubilizing the virus to provide soluble virus antigen having a molecular weight such that when supported on a particle, the sensitized particle remains mono-dispersed. In general, when using the rubella virus antigen for the sensitization of a particle, the soluble antigen does not have a molecular weight in excess of 125,000 , and most generally not in excess of 100,000, as determined by acrylamide gel electrophoresis.

As hereinabove indicated, the surfactant is employed in an amount which is sufficient to disrupt and solubilize the virus and which does not destroy the antiqenic characteristics thereof (too much detergent may destroy the antigenic characteristics). In general, the surfactant to virus weight ratio is an amount of from 0.2:1 to about 5:1, preferably from about 0.5:1 to 1:1. The selection of an optimum amount is deemed to be within the scope of those skilled in the art from the teachings herein.

temperature which does not denature the virus proteins, with such temperature generally not exceeding about 30°C with a temperature of from 20°C to 25°C being most convenient. Similarly, the pH is selected so as to maintain stability, with the pH being generally at 8.5, with the optimum pH generally being in the order of from 8.0 to about 9.0.

The treatment of the purified virus is effected at a

The treatment of the purified virus with the surfactant is for a period of time sufficient to disrupt the virus and effect solubilization thereof. In general, such disruption

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and solubilization can be accomplished in time periods in the order of from 5 to 120 minutes, however, in some cases longer or shorter times may be applicable.

The selection of an optimum treatment time is deemed to be within the scope of those skilled in the art from the teachings berein.

Applicant has found that by using a surfactant to disrupt
and solubilize the whole rubella virus, as hereinabove described,
it is possible to provide soluble rubella virus antigen which
tretains its antigenicity.

17 A procedure for disruption and solubilization of 12 whole virus, as hereinabove described, has been previously practiced in the art; for example, Vaheri et al. "Structural Proteins 13 14 and Subunits of Rubella Virus", Journal of Virology, P. 10-16 15 (Jan. 1972). In addition, it is known that such a procedure 16 is capable of recovering the structural proteins of the whole 17 rubells virus, with there being three principal structural 18 proteins, namely a structural protein with a molecular weight in 19 the order of from 60,000 to 65,000 daltons, a structural protein 20 with a molecular weight in the order of from 40,000 to 50,000 21 daltons, and a structural protein having a molecular weight in 22 the order of from 32,000 to 38,000 daltons. Applicant has also 23 found evidence of a structural protein having a molecular weight of from 100,000 to 120,000 daltons.

Applicant has found that the structural proteins recovered by such a procedure retain antigenic characteristics, and in addition, such structural proteins can be used in an assay for rubella antibody. Furthermore..., applicant has found that such structural proteins are capable of detecting early phase rubella.antibody, i.e., the rubella antibody present in serum or plasma within ten days of onset of rubella rash. The

term "rubella virus antigen" as used herein encompasses one or more of such structural proteins recovered bysuch

1 | procedure.

The hereinabove described technique for disruption and solubilization of whole rubella virus to provide soluble rubella virus antigen is also applicable to providing virus antigen from other viruses; e.g., those hereinafter disclosed with reference to a purification of virus. Such viral antigens may then be supported on a solid support, as hereinafter described, to provide a solid sensitized with the viral antigen for use in an assay.

In accordance with an aspect of the present invention, applicant has found that disruption and solubilization of whole rubella virus produces a soluble product which is antigenic and which is capable of reacting with rubella antibody, including the early phase antibody. Thus, by using a product prepared by such a procedure in an assay for rubella antibody; and in particular on a solid support, it is possible to detect rubells antibody even during the early phase.

As hereinafter described, the recovered product is of particular value for a direct agglutination assay, and applicant has found that such soluble rubella virus antigen may be supported on a latex particle (in particular a polystyrene) without the problem of self agglutination, i.e., the sensitized particles remain mono-dispersed.

The purified whole virus which is treated with surfacants is a virus which is produced in a tissue culture by procedures known in the art, and which is subsequently purified to remove non-virus lipids, nucleic acids, and non-viral proteins.

The tissue culture growth of rubella virus wherein rubella virus infected cells are raised in a suitable culture medium is well known in the art. The cells that are suitable for tissue culture growth to produce the rubella virus includes Vero cells, Baby Hamster Kidney, Procine Stabile Kidney, Serum

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Institute Rabbit Cornea and the like. In general, tissue cultures conventionally used for producing rubella virus are also suitable for the purposes of the present invention.

The virus may then be purified by procedures known in the art; e.g. as disclosed by Baheri et al., <u>supra</u>. In accordance with a preferred embodiment, the virus is purified in accordance with a procedure of the present invention.

More particularly, the procedure for purifying virus in accordance with the invention, involves, treating concentrated virus with hydroxyl apatite gel in an aqueous solution of controlled ionic strength and pH.

More particularly, after filtration and concentration, the virus is contacted with hydroxyl apatite gel in an aqueous solution having an ionic strength which is great enough to minimize or prevent adsorption of the virus by the gel, and which is low enough to allow the non-virus proteins to be adsorbed by the gel. The ionic strength is maintained by the use of phosphate ions, with the phosphate ions being present at a molarity of from 0.05M to 1.5M to provide for effective adsorption of non-virus proteins and nucleic acids, without significant adsorption of the virus. The phosphate molarity in most cases is at least 0.08 M.

In addition, the adsorption is conducted at a pH in the order of from 6 to 9, most generally in the order of from 7 to 8. The pH of the solution is maintained by the use of a suitable buffer. The adsorption may be conducted in the presence of EDTA at a concentration from .OlM to .O001M. EDTA as well as other chelating agents increases adsorption of non-viral proteins and nucleic acids, and aids in minimizing the adsorption of viral proteins.

By proceeding in accordance with the purification of the invention, the high molecular weight proteins and nucleic acids are adsorbed by the gel to thereby separate the virus protein from the non-virus proteins having similar molecular weights.

After such adsorption, the lower molecular weight proteins 2 still remaining in the fluid may be separated by conventional procedures. Thus, for example, further separation may be accomplished by centrifugation through a barrier layer or cushion as known in the art. In particular, the virus protein is centrifuged through a suitable barrier layer such as sucrose. glycerol, cesium chloride, cesium sulfate and the like, with the lower molecular weight proteins remaining above the barrier, and the virus being centrifuged through the 10 barrier, as a separate layer. The fluid containing the low molecular weight proteins and the barrier layer are then removed leaving a virus protein essentially free of non-virus proteins, nucleic acids, lipids, and the like. In general, the purified virus contains less than 1%, most generally less than 0.1% of non-virus lipids, nucleic acids and proteins. 16

The above procedure may be employed for purifying a wide variety of viruses, including, but not limited to: rubella virus; rubeola virus, herpeto viruses (herpes simplex ,Varicella Zoster, cytomegalovirus, Epstein-Barr [infectious mono-nucleosis]) parainfluenza viruses; influenza virus; dengue virus, etc.

Such purified virus may then be treated with a surfactant to disrupt the virus and effect solubilization thereof to thereby provide a virus antigen, as hereinabove described.

It is to be understood that although the hereinabove described procedure for purifying the rubella virus is preferred, other procedures for separating non-virus proteins, lipids and nucleic acids can also be employed for purifying the rubella virus for subsequent treatment with surfactant to thereby produce the soluble rubella virus.

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The viral antigen which is prepared by disruption and solubilization of whole virus may be supported on a solid support for use in an assay. The following description is particularly directed to rubella virus antigens; however, the teachings are also applicable to other viral antigens.

The rubella virus antigen prepared by disrupting and solubilizing whole rubella virus may then be supported on a solid support for use in an assay for rubella virus antibody. Such supported rubella virus antigen is capable of reacting with early phase rubella virus antibody. In accordance with the preferred embodiment, the rubella virus antigen is supported on a particula support for use in an agglutination assay; however, it is to be understood that the rubella virus antigen may be supported on a non-particulate support (or for that matter on a particulate support) for use in an assay for rubella virus antibody by procedures other than the agglutination technique. Thus, for example, the supported rubella virus antigen may be supported on a solid support for use in an assay for rubella virus antibody by a radioimmunoassay, fluorescent or enzyme assay technique. Similarly, the rubella virus antigen of the present invention may be employed for the assay of rubella virus antibody in unsupported form by use of such techniques. Thus, the scope of the invention is not limited to the preferred embodiment wherein the rubella virus antigen is supported on a particulate support for use in an agglutination assay for rubella virus antibody.

The antigen may be supported on any one of a wide variety of solid supports which are capable of supporting the antigen, and which can be used in the assay procedure without interfering with the immunochemical reaction. Moreover, the support should be one which is stable; i.e., not adversely affected by the prepared antigen. The antigen may be supported on the support by an adsorption technique, or by covalent coupling,

either by activation of the support, or by the use of a suitable coupling agent, or by use of reactive groups on the support.

Such procedures are generally known in the art.

The support may be any one of a wide variety of supports,
and as representative examples of suitable supports there may
be mentioned: synthetic polymer suppports, such as polystyrene,
polypropylene, substituted polystyrene (e.g., aminated or
carboxylated polystyrene), polyacrylamides, polyamides, polyvinylchloride, etc.; glass beads, agarose; etc. The supports
may include reactive groups; e.g., carboxyl groups, amino groups
etc. to permit direct linking of the virus antigen to the
support.

In accordance with preferred emodiment, the particulate support is either a polystyrene, amivated polystyrene, carboxylated polysytrene or a polyvinylchloride, although, it is to be understood that the scope of the invention is not limited to such supports.

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As hereinabove indicated, the antigen may be supported on the support by the use of an adsorption technique, or by covalent coupling with a coupling agent. As representative examples of suitable coupling agents there may be mentioned: dialdehydes; for example glutaraldehyde, succinaldehyde, malonaldehyde, etc; unsatured aldehyde, e.g., acrolein, methacrolein, crotonaldehyde, etc.; carbodiinides; diisocyanates; dimethyladipimate; cyanuric chloride etc. The selection of a suitable coupling agent should be apparent to those skilled in the art from the teachings herein.

27 Similarly, the antigen may be supported by activation
28 of a suitable support; for example, cyanogen, bromide activated
29 magazose.

In accordance with a preferred embodiment, as hereinabove

noted, the soluble rubella virus antigen is supported on a particulate support which is either polystyrene (substituted or unsubstituted) or polyvinylchloride; most preferably polystyrene.

In some cases, the soluble antigen may be supported by an adsorption technique, in other cases, it may be necessary to employ covalent coupling.

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The virus antigen sensitized particulate support is preferably prepared for use in an assay in which rubella virus antibody is determined by an agglutination technique. The particulate support is provided with an effective amount of the antigen for the assay, while preventing excessive amounts which may result in bridging of the antibody to a single particle. In general the weight ratio of soluble rubella antigen to support is from 1:100 to 1:5000. The selection of an optimum amount is deemed to be within the scope of those skilled in the art from the teachings herein.

17 In accordance with one technique, after the antigen is adsorbed on the particles, the support, including the adsorbed antigen, is further coated with protein which does not 20 adversely affect the subsequent immunochemical reaction in order to provide a protein coating on the portion of the support which does not including the antigen. As should be apparent, 23 the protein coating should not immunologically 24 react with either the rubella virus antigen or with sera 25 to be used in the assay. As examples of suitable proteins there 26 may be mentioned: bovine serum albumin, ovalbumin, and the like. 27 The selection of a suitable protein to saturate the spaces 28 between the rubella virus antigen on the support is deemed to 29 be within the scope of those skilled in the art from the teachings herein.

It is to be understood that such coating with protein
is not required for producing sensitized particles for use in
an agglutination assay.

After the rubella virus antigen has been supported on a solid support, as generally practiced in the art for the production of sensitized particles for use in an agglutination assay, the sensitized particles are treated with a liquid containing only many hylene sorbitan monolaurate (Tween 20) at a weight ratio to the polystyrene of 0.1.1 to 1

The sensitized particles are preferably a synthetic polymer and in particular a polystyrene [substituted (carboxylated or aminated) or unsubstituted) or polyvinylchloride latex. Applicant has found that sensitization of such particles with soluble rubella virus antigen prepared, as hereinabove described, produces a sensitized particles which remains mono-dispersed (no self agglutination), whereby such sensitized latex particles may be effectively employed in a direct agglutination assay for rubella antibody. Such sensitized particles are capable of detecting early phase rubella antibody. In addition, such sensitized particles are capable of providing a direct agglutination assay having a high sensitivity for rubella antibody.

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The rubella virus antigen sensitized particle prepared in 21 accordance with the invention are suitable for use in a kit and assay for rubella virus antibody by a direct agglutination 23 procedure. Such kit may include, in addition to the sensitized 24 rubella virus particles, as hereinabove described, in a suitable 25 container therefor, a reactive serum control (contains rubella 26 antibody) and a non-reactive serum control (no rubella antibody) 27 in suitable containers therefor. In accordance with a preferred 28 embodiment, in addition to the reagents, there is provided a 29 test card on which the assay is effected. The test card has a 30 31 | flat testing surface which include suitably marked areas (for 32 example, a test circle) for placing one or more samples to be assayed, as well as suitably marked areas for each of the serum

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controls. The test card and reagents may be included in a single kit package.

In the aggultination assay, undiluted serum or diluted serum (e.g. 1:10) is contacted with the sensitized particles followed by mixing, with the presence of the antibody against rubella virus being evidenced by visible agglutination.

Such rubella virus antigen sensitized particles may also be employed in a quantitative assay for rubella virus antibody.

9 In a quantitative assay, the sample to be assayed is
10 serially diluted, as appropriate, and to each serial dilution
11 there is added the particles sensitized with the soluble rubella
12 antigen. The quantity of antibody in the sample is determined from
13 the highest dilution giving any agglutination of the sensitized
14 particles.

The quantitative or qualitative assay for rubella
antibody may be effected on a card surface wherein the surface
includes suitably marked areas for placing the sample and
control to which the senitized particles are added.

The invention will be further described with respect to the following examples; however, the scope of the invention is not to be limited thereby:

EXAMPLE I

Production and Purification of Rubella Virus.

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Confluent roller cultures (680 cm²) of Vero cells (a
continuous culture line of cells derived from African Green monkey
kifney) were innoculated with approximately 0.01 PFU of rubella virus

reper cell and maintained in a standard culture medium (Medium
leg 199) containing .025 M hepes buffer, pH 7.4, and 2% (vol/vol) of
the filtrate obtained by forcing fetal bovine serum through a
membrane designed to retain molecules of 100,000 molecular

weight and greater (Amicon XM-100 membrane). The medium was changed daily, and the culture fluids having a hemagglutination titer greater than 16 were made to contain 0.01 M Tris base and 0.01 M EDTA. After incubation at 4°C for 1 hour, they were concentrated . in an Amicon hollow fiber dialyzer-concentrator to 1/10 the original volume. After clarification at 5,000 x g for 20 minutes, the pH was adjusted to 7.6 at 22°C and 1/10 volume of hydroxylapatite suspension was added, and the slurry was incubated at 4°C with mixing, overnight. The hydroxylapatite was removed by centriquation at 5,000 x g for 15 minutes, after which 30 ml of the concentrate was layered over 9 ml of 69% (wt/wt) glycerol in a Beckman SW28 tube. The virus was sedimented at 82,000 x g for 16 hours at 4°C, and the resultant pellet was resuspended in 0.01 M carbonate buffer, pH 9.5 (coating buffer). The purified virus was assayed for hemagglutinin content and stored at -70°C.

EXAMPLE II

Solubilization of Purified Virus.

The purified virus in 0.01 M carbonate buffer, pH 9.5, was solubilized by treatment with sodium dodecyl sulfate (SDS). The purified virus was made to contain 0.05% (w/v) SDS and was incubated for 30 minutes at room temperature.

EXAMPLE III

Preparation of Sensitized Latex.

Commercial suspensions of polystyrene latex (0.9 micron diameter particles) were washed four times with 25 volumes each of the coating buffer and were resuspended in the coating buffer to provide 3% solids (Vol/vol). The latex suspension was added directly to the solubilized virus at a ratio of 2 volumes of the

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3% latex to 1 volume of solubilized virus and the suspension was mixed by tumbling for 16 hours at room temperature. The sensitized latex was washed twice with 20 volumes of 1% bovine serum albumin in phosphate buffered saline (BSA-PBS) and resuspended at 0.5% in 1% BSA-PBS contained 0.05% polyoxyethylene sorbitan monolaurate surface active agent (Tween 20) and 0.02% gentamiacin.

EXAMPLE IV

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Latex Agglutination Test for Rubella Virus Antibodies. Glass plates with 1.4 cm fused circles were employed. Serial 2-fold dilutions of serum were prepared in 1% BSA-PBS-Tween 20 and 25 ul of each dilution was placed in separate wells. After adding 25 ul of sensitized latex, the serum and latex suspension was mixed and rotated 100 rpm for 5 minutes. The presence of antibody against rubella virus was evidenced by 15 visible agglutination.

EXAMPLE V

17 Purified virus prepared in accordance with Example I was treated with a 1% aqueous solution of sarcosyl for 30 minutes 18 at room temperature in coating buffer to disrupt and solubilize 19 the virus.

21 The pH of the solubilized virus was adjusted to 6.5 with hydrochloric acid and mixed with two volumes of 3% carboxylated 22 polystyrene latex (in phosphate buffer, pH 6.5) for 1 hour at 23 24 4°C.

25 To the solution was added 10 mg of a carbodiimide 26 coupling agent and the mixture was mixed overnight at 4°C.

After centrifugation, the solids were resuspened in phosphate buffered saline (PBS) followed by centriquation and resuspension in PBS containing 1% BSA and 0.05% Tween 20.

The procedure covalently bound the soluble rubella virus antigen to the latex.

EXAMPLE VI

In accordance with a preferred procedure, there is provided a test card for rubella antibody. The test card includes a marked circle for a reactive control, a marked circle for non-reactive control, as well as one or more test sample circles.

25 ul of undiluted serum sample is placed in an appropriately marked sample circle, and 25 ul of the reactive and nonreactive controls are placed in their respective circles.

with a micropipettor, there is added sensitized latex of Example III (approximately 15 ul), followed by rotation on a rotator (about 8 minutes), and gentle hand rotation.

The card is read microscopically in the wet state under lamp a high intensity incandescent lamp.

14 The reactive control should show definite aggultination 15 and the non-reactive control should show no agglutination.

16 Any serum samples showing any agglutination should be re17 ported as reactive.

Numerous modifications and variations of the present invention are possible in light of the above teachings and, therefore, within the scope of the appended claims, the invention may be practised otherwise than as particularly described.

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WHAT IS CLAIMED IS:

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- A composition, comprising:
- a solid support sensitized with soluble rubella virus antigen, said soluble rubella virus antigen having been derived by disruption and solubilization of whole rubella virus.
- The composition of Claim 1 wherein the solid support is a particulate support.
- 9 3. The composition of Claim 2 wherein the soluble rubella
 10 virus antigen has a molecular weight of no greater than 125,000
 11 daltons as determined by acrylamide gel electrophoresis.
 - The composition of Claim 3 wherein the particulate support is a polystyrene latex.
 - 5. The composition of Claim 1 wherein the antigen on the support is comprised of at least one of the structural proteins of the virus and the supported antigen is immunoreactive with early phase rubella antibody.
 - The composition of Claim 5 wherein the antigen is supported on a particulate support.
 - The composition of Claim 6 wherein the sensitized solid particles remain monodispersed.
 - The composition of Claim 7 wherein the solid particle is a polystyrene latex.
 - The composition of Claim 3 wherein the solid support is a synthetic polymer.
 - 10. The composition of Claim 9 wherein the synthetic polymer is selected from the group consisting of polyvinylchloride, polystyrene, aminated polystyrene and carboxylated polystyrene.
 - 11. The composition of Claim 9 wherein the antigen is covalently coupled to the solid support.

12. The composition of Claim 9 wherein the antigen is adsorbed on the solid support.

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13. The composition of Claim 2 wherein the disruption and solubilization of the whole rubella virus is effected with a detergent.

14. The composition of Claim 13 wherein the detergent is an alkali dodecyl sulfate.

15. A process for producing a solid support sensitized with soluble rubella virus antigen, comprising:

deriving soluble rubella virus antigen by disruption and solubilization of whole rubella virus, and supporting the soluble rubella virus antigen on a solid support.

- 16. The process of Claim 15 wherein the solid support is a particulate support.
- 17. The process of Claim 16 wherein the soluble rubella virus antigen has a molecular weight of no greater than 125,000 daltons, as measured by acrylamide gel electrophoresis.
- 18. The process of Claim 17 wherein the particulate support is a polystyrene latex.
 - The process of Claim 15 wherein the supported antigen is comprised of at least one of the structural proteins of the virus and is immunoreactive with early phase rubella antibody.
- 20. The process of Claim 19 wherein the antigen is supported on a particulate support.
- 21. The process of Claim 20 wherein the sensitized solid particles remain monodispersed.
- 22. The process of Claim 21 wherein the solid particle is a polystyrene latex.
- 23. The process of Claim 17 wherein the solid support is a synthetic 'polymer.

- 25. The process of Claim 23 wherein the antigen is covalently coupled to the solid support.
- 26. The process of Claim 23 wherein the antigen is adsorbed on the solid support.
- 27. The process of Claim 16 wherein the disruption and solubilization of the whole rubella virus is effected with a detergent.
- 28. The process of Claim 27 wherein the detergent is an alkali dodecyl sulfate.
 - 29. In a kit for determining rubella virus antibody by an agglutination technique, the improvement comprising:
- said kit including in a reagent container solid particles
 sensitized with soluble rubella virus antigen, said soluble
 rubella virus antigen having been derived by disruption and
 solubilization of whole rubella virus.
 - 30. The kit of Claim 29 wherein said kit further includes a test card having a flat surface for receiving assay samples.

 - 32. The kit of Claim 30 and further comprising in separate reagent containers a reactive serum control of rubella antibody and a non-reactive serum control free of rubella antibody.
 - 33. In a direct agglutination assay for rubella virus antibody employing solid particles sensitized with rubella virus antigen, the improvement comprising:
 - said rubella virus antigen having been derived by disruption and solubilization of whole rubella virus.

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34. The assay of Claim 33 wherein the soluble rubella virus antigen has a molecular weight of no greater than 125,000 daltons as determined by acrylamide gel electrophoresis.

- 35. The assay of Claim 34 wherein the particles are a polystyrene latex.
- 36. The assay of Claim 35 wherein the sensitized particles are comprised of at least one of the structural proteins of the virus and are immunoreactive with early phase rubella antibody.
- 37. The assay of Claim 34 wherein the particles are a synthetic polymer.
- 38. The assay of Claim 37 wherein the synthetic polymer is selected from the group consisting of polyvinylchloride, polystyrene, aminated polystyrene and carboxylated polystyrene.
- 39. The assay of Claim 37 wherein the antigen is adsorbed on the particles.
- 40. The assay of Claim 37 wherein the antigen is covalently coupled to the particles.
- The assay of Claim 34 wherein the disruption and solubilization of the whole rubella virus is effected with a detergent.
- 42. The assay of Claim 41 wherein the detergent is an alkali dodecyl sulfate.
- 43. In a process for purifying whole virus to separate the virus from non-viral proteins, the improvement comprising:
- contacting the whole virus with hydroxyl apatite in the presence of phosphate ion and at a pH of from 6 to 9, said phosphate ion being present in a molarity of from 0.05 M to 1.5M to provide for adsorption of non-viral proteins without significant adsorption of viral protein.
- 44. The process of Claim 43 wherein the virus is rubella virus.

5. The process of Claim 44 wherein the pH is from 7 to 8.

46. A composition, comprising:

a solid support sensitized with viral antigen, said viral antigen having been derived by disruption and solubilization of whole virus.

47. In an assay for rubella virus antibody wherein rubella virus antibody immunoreacts with rubella virus antigen, the improvement comprising:

immunoreacting in said assay rubella virus antibody with soluble rubella virus antigen derived by disruption and solubilization of whole rubella virus.

EUROPEAN SEARCH REPORT

84 10 5636 EP

DOCUMENTS CONSIDERED TO BE RELEVANT				CLASSIFICATION OF THE
Category	Citation of document with i	ndication, where appropriate, t passages	Relevant to claim	APPLICATION (Int. Cl.4)
Y	CHEMICAL ABSTRACT no. 3, January 22 256, abstract 14: OHIO, US); A.A. 5: "Characterization structural antige virus reacting precipitation", 6: Microbiol. Scand 80(4), 534-44 * abstract *	2, 1973, page 339q, (COLUMBUS, SALMI: 1 of a en of rubella 2 gel 3 Acta Pathol.	1-47	G 01 N 33/56 G 01 N 33/54 C 12 N 7/06 C 12 N 7/02
¥	CHEMICAL ABSTRAC no. 1, July 4, 1 abstract 2755t, US); M.N. WAXHAM "Immunochemical of rubella virus & Virology 1983, * abstract *	983, page 275, (COLUMBUS, OHIO, et al.: identification	1-47	TECHNICAL FIELDS SEARCHED (Int. CI.4)
D,A	US-A-4 195 074 * the entire doc	- (J. SAFFORD Jr.) ument *	1	G 01 N C 12 N A 61 K
Y	GB-A-2 001 326 * abstract; line	(SANDOZ LTD.)	1-47	
Y	EP-A-0 054 249 INDUSTRIES INC.) * the entire doc		1-47	
		/-		
-	The present search report has t	oeen drawn up for all claims		
	Place of search THE HAGUE	Date of completion of the search	OSBO	DRNE H.H.

CATEGORY OF CITED DOCUMENTS

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- T: theory or principle underlying the invention
 earlier patent document, but published on, or after the filling date
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EUROPEAN SEARCH REPORT

016,1328

EP 84 10 5636

	DOCUMENTS CONSI	Page 2		
ategory	Citation of document with of releva	indication, where appropriate, nt passages	Relevant to claim	CLASSIFICATION OF THE APPLICATION (Int. Cl.4)
Y	US-A-3 622 663 * abstract; clai		43,44	
A	EP-A-0 001 838 AKTIENGESELLSCHA	- (BEHRINGWERKE FT)		
A	CHEMICAL ABSTRAC no. 1, January 6 abstract 1147k, US); A.E. AULETT "Effect of sodiu on rubella virus Microbiol. 1968,	, 1969, page 98, (COLUMBUS, OHIO, A et al.:		
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				TECHNICAL: FIELDS SEARCHED (Int. CI.4)
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	Ŧ			
	The present search report has t	and drawn in for all claims	-	
			ــــــــــــــــــــــــــــــــــــ	Examiner
	THE HAGUE	Date of completion of the search 18-01-1985	OSBO	RNE H.H.
Y : p.	CATEGORY OF CITED DOCI articularly relevant if taken alone articularly relevant if combined w ocument of the same category	E : earlier pa after the f with another D : documen	tent document	
0 : n	echnological background on-written disclosure ntermediate document	& : member of document		tent family, corresponding